



Hydrodeoxygenation of acetone–furfural condensation adducts over alumina-supported noble metal catalysts



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ABSTRACT

The hydrodeoxygenation (HDO) of aldol condensation adducts is of key interest in the preparation of biofuels from biomass by catalytic routes. In this work, furfuryldeneacetone is taken as model of biomass-derived condensation adduct with furanic rings, unsaturations and carbonyl functionalities. Four noble metal catalysts (alumina-supported Ru, Rh, Pd and Pt) were tested. Obtained results show that Rh and Ru catalysts are only active for the hydrogenation of aliphatic double bonds; whereas Pd, and specially Pt catalysts, are active for the total HDO of the adduct, yielding *n*-octane, with selectivities higher than 30% at 493 K after 24 h on stream (total conversion and carbon balance closure higher than 90%). Based on these results, a mechanism (considering serial, parallel and equilibrium steps) and the corresponding kinetic model have been proposed for explaining the different selectivity trends observed for these metals.

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1. Introduction

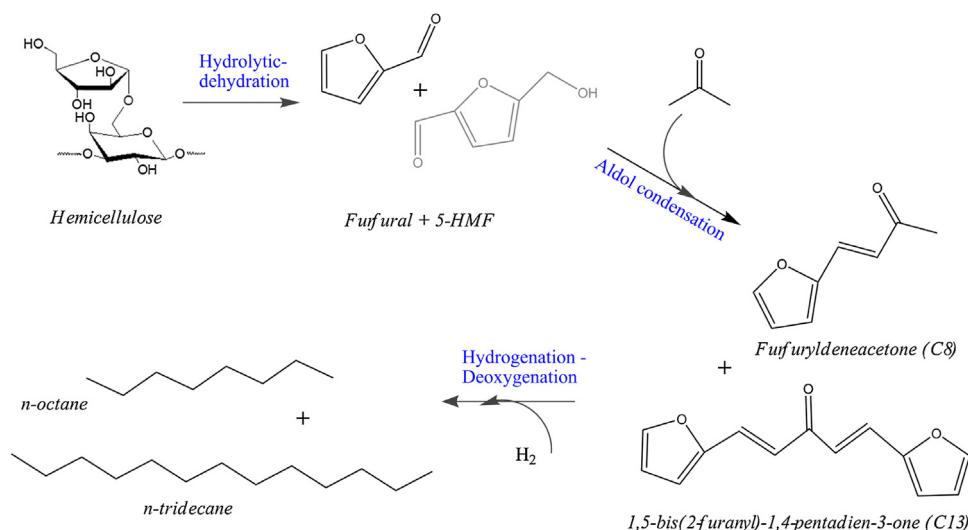
Biomass constitutes a promising raw material for renewable fuels and chemicals, especially for the manufacture of drop-in liquid fuels [1]. Aldol condensations of sugar-derived aldehydes are aqueous phase processes with significant advantages over thermochemical and biological methods [2]. These reactions use as raw materials a large amount of bio-based platform molecules, and can be performed at ambient conditions [3,4]. Hydrodeoxygenation (HDO) of the condensation adducts leads to drop-in fuels, (from C8 to C15 lineal alkanes, a diesel-quality fraction) [5,6]. Gasoline-range hydrocarbons can also be obtained from alcohols, aldehydes or ketones, products of the fast pyrolysis of the lignocellulosic biomass through condensation-hydrogenation sequences [7,8].

Hydrodeoxygenation of these condensation adducts to *n*-alkanes requires several steps to remove the oxygen atoms present in different functional groups (C=O, C–OH, C–O–C): hydrogenation of aliphatic insaturations (C=C), furan-ring hydrogenations, heterocyclic ring opening and dehydration/hydrogenation steps. The complexity of this reaction is largely increased due to the steric interactions of furan rings, which modify the general mechanism of individual reactions [9].

A limited number of works devoted the total HDO of these compounds have been published. Contrary, the hydrogenation of furanyl compounds to alcohols has been deeply studied in gas phase. High selectivities with Co and Ni-Fe catalysts at temperatures higher than 500 K have been reported, although with a strong catalytic deactivation [10,11]. Less deactivation was observed in liquid phase HDO over Ni supported catalysts [12,13], with selectivities higher than 98%. Likewise, dehydration/hydrogenation processes to remove oxygenated groups were deeply studied for functionalized lineal hydrocarbons over noble metals (Pt group) and sulfide catalysts [14]. The sulfur leaching, as well as the formation of undesirable S-containing products, prevents the use of sulfide catalysts when using biomass-derived compounds [15]. In addition, the concentration of sulfur compounds is lower in the case of streams coming from biomass processes than in petrochemical processes. It should be also noted that in lignocellulose materials, sulfur is mainly associated to the lignin, and therefore is removed in the pretreatment of these materials. Consequently, noble metals seem to be the optimum catalysts for these reactions. Dehydration was also observed as side-reaction in the hydrogenation of furfural using Cu/SiO₂ at temperatures higher than 523 K [16] and in hydrogenations of polyalcohols in presence of Pt or Ru and acid sites [17]. In presence of Pd, competition of this reaction with decarbonylation was observed [18].

Furmisky et al. reviewed the application of HDO reactions to small aromatic compounds obtained from biomass, using oxides supported on alumina as catalysts [19]. More recent works

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Scheme 1. General mechanism for biodiesel synthesis from hemicellulosic biomass by dehydration-aldolization-hydrogenation/deoxygenation.

dealing with the biomass derived fraction (instead of the oil-derived one) used Pt-based catalysts at 723 K and 5 MPa [20], or Pd-based catalysts working at supercritical conditions [21]. Different side-products and hydrogenated intermediates were detected, limiting the selectivity of corresponding alkanes. Likewise, it was observed that lower temperatures and pressures are desirable to prevent coking and side reactions, thus the convenience of carrying out the reaction in liquid phase is clear [22]. However, there is no agreement in the hydrodeoxygenation mechanism. Therefore, this process has been usually presented for sugar-derived aldehydes as two independent reactions: a first hydrogenation of condensed adducts removing the aliphatic C=C insaturations (catalyzed by noble metals at moderate temperature and pressure) [23] and, a second step of four dehydration/hydrogenation (4ADP/H) reactions at stronger conditions (using bifunctional catalyst with metal and acid sites) which produce the alkanes [20,24].

In this work, the total hydrodeoxygenation of condensed adducts in one step is studied. This reaction is the last step in the manufacture of diesel blends from cellulosic biomass (Scheme 1). The main scope of this work is to identify the most effective active phase, the optimum reaction conditions and to propose reaction pathways for the hydrodeoxygenation of condensed adducts. The study is focused on the furfuryldeneacetone (C8), the first condensation adduct obtained by aldolization of furfural and acetone [4]. Four different noble metals (Ru, Rh, Pd and Pt) supported on alumina were tested at different temperatures. Group VI–VIII metals have been previously tested in hydrogenations at low-medium temperatures showing high catalytic activity, but also a high capacity of adsorption and activity for side reactions [25]. Pd and Rh are considered good catalysts for hydrogenation of aliphatic insaturations [26,27] whereas Ru and Pt show high activity in deoxygenation steps [28,29]. Alumina was chosen as support because of its acidity, which can favor some steps of the HDO reaction, but it is considered mild enough for not promoting coke formation reactions.

2. Experimental

2.1. Catalysts characterization

Catalysts used in this work are BASF Escat series of 0.5% (w/w) of Pd, Rh, Ru or Pt supported on γ -Al₂O₃. All the catalysts were activated at 473 K for 2 h under flowing hydrogen. Different studies confirm that these metals are completely active after this

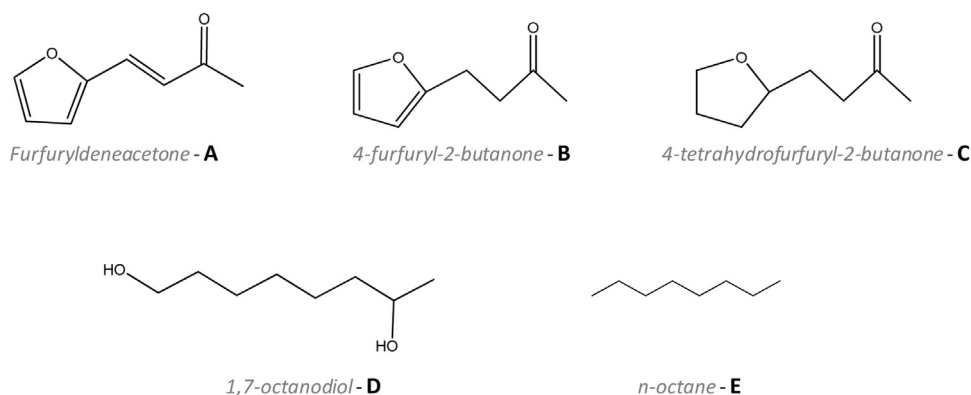
pretreatment (the position of the metal on the surface favors the C=O hydrogenation), whereas higher reduction temperatures promote active phase sintering and the formation of different structures on the surface catalytically active for other side reactions [30].

Morphological properties were determined in previous works by using the nitrogen physisorption at 77 K in a Micromeritics ASAP 2020 analyzer by the BET method for the specific surface area and the BJH approach to determine the pore volume and diameter [31,32]. The metal particle morphology and size distribution were identified by TEM using a JEOL-2000 EX-II microscope. The strength and distribution of the acid sites were determined by temperature programmed desorption (TPD) of pre-adsorbed NH₃ in a Micromeritics TPD/TPR 2900. Samples (10 mg) were treated in He at 623 K for 2 h and exposed to a 2.5% NH₃ in He stream at 323 K until saturation. After the stabilization, the temperature was increased at a rate of 5 K/min and the NH₃ evolution was monitored by mass spectrometry. Further details about the acidity measurements are given in previous works [33].

2.2. Reaction studies

Reactions were carried out in a 0.5 L stirred batch autoclave reactor (Autoclave Engineers EZE Seal) equipped with a PID temperature controller and a back pressure regulator. The reactor was loaded with 0.25 L of a hexane solution of 0.42 g of furfuryldeneacetone (Alfa Aesar, 98%). The catalyst (60 mg), with an average particle diameter of 50–80 μ m, was added and air is purged out by adding nitrogen up to 1.5 MPa for three times before starting the reaction. The concentration of C8, as well as the catalyst/reactant ratio was chosen considering the C8 maximum concentration obtained in previous works related to the furfural–acetone aldolcondensation [4]. Selected reactant concentration also ensures that the solubility of the reactants or reaction products does not affect catalytic performance.

The reactor was initially pressurized to 2.5 MPa with H₂, heated to reaction temperature (reaching final pressures of 5.5 MPa) and stirred at 1000 rpm for 8 h for the screening of the different metals (reactions at 393 K) or 24 h in the studies of the influence of the temperature (393, 433, 453 and 493 K). Selected pressure guarantees the H₂ excess during all the experiment, since the amount of hydrogen needed for the total hydrogenation of the adduct was less than a 10% of the total amount of hydrogen loaded in the reactor. This fact was also experimentally checked both experimentally, the pressure after 24 h only decreases in 0.3–0.5 MPa when hydrogen



Scheme 2. Chemical structure of the reactant (A) and reaction products identified by GC-MS during the hydrodeoxygenation of furfuryldeneacetone.

is not fed; and theoretically, assuming the Henry's equilibrium for the hydrogen-hexane system.

Samples were withdrawn from the sampling port during the reaction, filtered and analyzed by capillary GC in a Shimadzu GC-2010 equipped with a FID detector, with a CP Sil 5 capillary column as stationary phase. Peak assignment was performed by GC-MS and responses were determined using standard calibration mixtures and the effective carbon number concept [34]. The structure of the different isomers can be observed in Scheme 2. Due to the complexity of their nomenclature, each compound was labeled with a capital letter that will be used during the discussion.

The stability of the catalysts was tested by repeating the batch experiments with recovered catalysts. The maximum decrease in conversion observed was 10%, whereas the selectivities of total hydrogenated products showed just slight decreases (<5%). Therefore, catalyst deactivation has not being considered in further discussions. Characterization of the used catalysts (nitrogen physisorption, XRD) does not reveal any morphological or crystallographic change during the reaction.

3. Results and discussion

3.1. Catalytic activity

Batch experiments at 393 K and 4.5 MPa were performed with the selected catalysts in order to identify the most active metal and determine the reaction pathways for the hydrogenation of the studied condensation adduct. This temperature was chosen considering previous works about the hydrogenation of other biomass-derived compounds [23]. The reaction time was in all cases 8 h. The most active catalyst was identified considering both the selectivity of n-octane (labeled as "E") and the carbon balance. The selectivity was calculated as the ratio between the concentration of n-octane and the concentration of converted C8-condensation adduct.

Reaction profiles obtained with the four catalysts are given in Fig. 1. These profiles are characterized by a fast disappearance of the reactant with total reactant conversion after less than 4 h (best results for Pt/Al₂O₃, with total conversion after 2 h). This drop in the reactant concentration is directly related to the appearance of

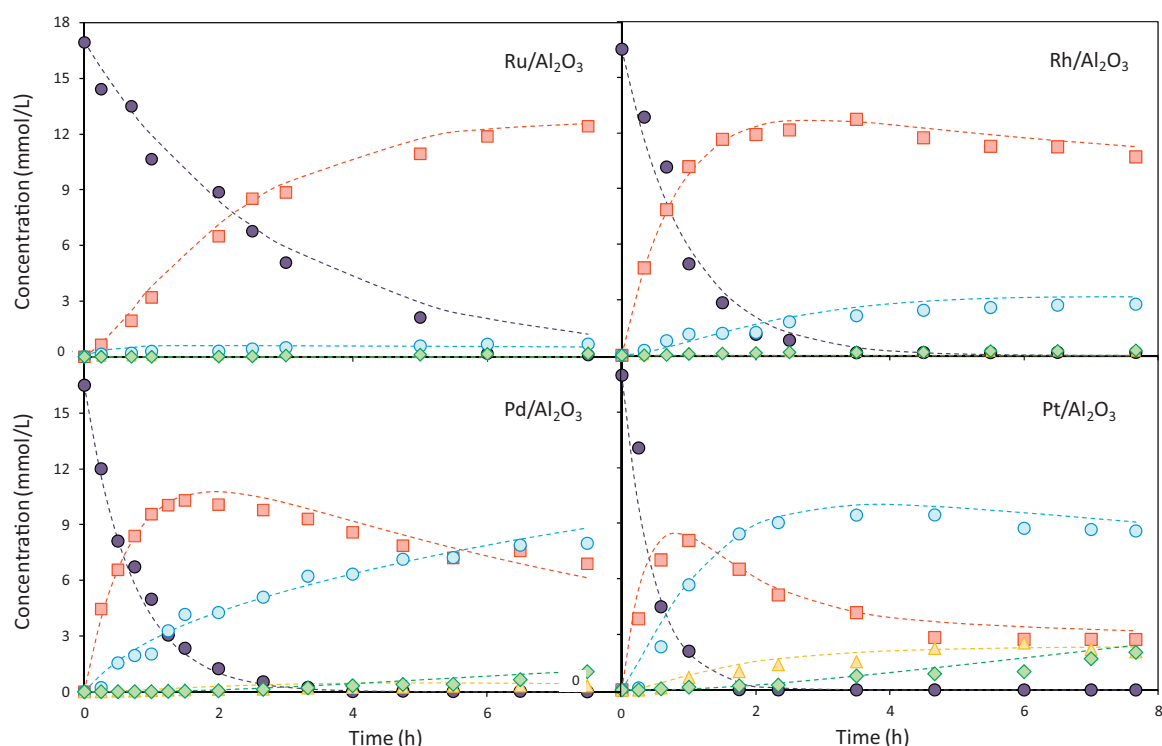


Fig. 1. Concentration profiles obtained in HDO of C8-condensate at 393 K. Symbols: (●) "A"; (■) "B"; (●) "C"; (▲) "D"; (◇) "E" (see Scheme 2 for notation).

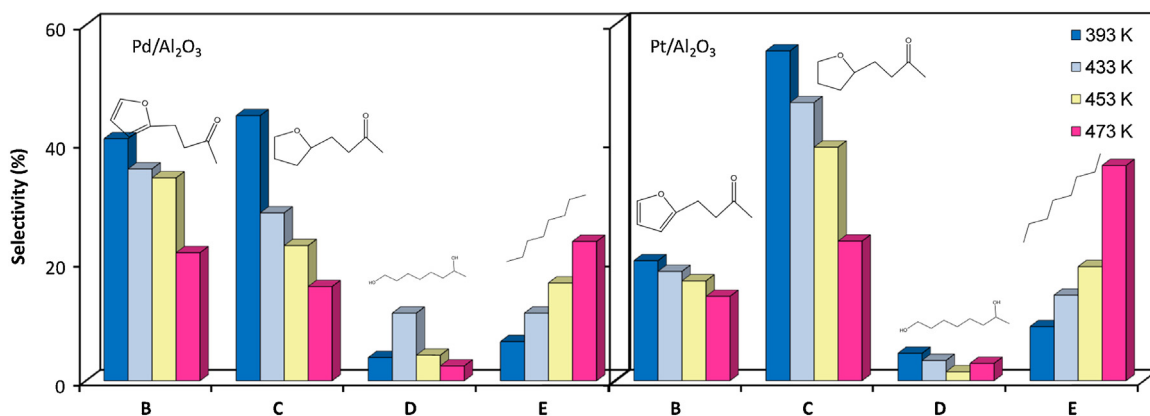


Fig. 2. Final selectivities obtained after 24 h of reaction at different temperatures using Pd/Al₂O₃ or Pt/Al₂O₃ as catalyst. Results obtained at 95% of conversion.

the first intermediate (labeled as “B”), characterized by the selective hydrogenation of the aliphatic C–C double bond. In the case of Ru and Rh supported catalysts, the profile of the B intermediate reaches a maximum (selectivities of 80–85%) and the concentration remains in values close to the maximum. On the other hand, when Pd or Pt/alumina are used as catalysts, the maximum of this intermediate is reached faster (less than 2 h) and its concentration drops to values close to zero at the end of the experiments. This decrease is more sharply for Pt/alumina due to the hydrogenation of more stable in unsaturations (cyclic-aromatic bonds, carbonyl groups) to obtain other intermediates (labeled as “C” and “D”) and the n-alkane (“E”). The most unstable intermediate is the open-cycle “D”, whose selectivity never reached values higher than 3%, although a maximum is clearly observed. Previous results confirmed that Pd is very selective for hydrogenation aliphatic unsaturated aldehydes [30], but Pt is much more active in deoxygenation reactions [29]. Consequently, the selectivity of n-octane obtained with Pt/alumina (9.4%) was higher than for Pd/alumina (6.1%). The carbon balance closures obtained were, in all the cases, higher than 95%, discarding the presence of other side reactions such as decarbonylations or the formation of heavy adsorbed by-products.

The modest n-octane selectivities obtained in both cases (with Pd and Pt) are due to the reaction parameters: the short reaction time (profiles suggest increasing trends for n-octane selectivity at longer times) or the reaction temperature. In fact, the complete hydrodeoxygenation of polyfunctionalized molecules was carried out at higher temperatures (over 650 K) to guarantee the complete reaction, with negligible activity at temperatures lower than 400 K [7]. The evolution of the activity at higher temperatures (433, 453 and 493 K) was tested with the most active catalysts (Pd and Pt/alumina). Fig. 2 compares selectivities after 24 h of reaction. In both cases, the n-octane selectivity increases with the temperature, reaching values of 25 and 35% with Pd and Pt, respectively. However, the differences between both materials are more noticeable analyzing the distribution of the different reaction intermediates. The first intermediate (“B”) reaches almost 40% of the final selectivity (at low and medium temperatures) on the Pd/alumina catalysts, whereas lower than 25% on Pt/alumina. The high selectivity of Pd for aliphatic unsaturated hydrogenations in contrast with the low activity in presence of cyclic compounds was previously observed in hydrogenations of compounds as benzaldehyde or acetophenone [30].

Concerning to the carbon balances, in both cases, decrease with the temperature. This behavior is more noticeable with Pd/alumina (from 97.7% at 393 K to 63.7% at 493 K), whereas for Pt/alumina the variation was lower than 10% (from 98.1% at 393 K to 88.8% at 493 K). At first insight, this carbon unbalance can be caused by side reactions leading to condensation products, such as Diels–Alder

reactions. At this point, the larger selectivities observed for the intermediate “B” with two conjugated C–C double bonds (active for Diels–Alder reactions) can be the responsible of these differences.

3.2. Kinetic study

The evolution of the selectivity of different products as function of the C8-condensated conversion is plotted in Fig. 3 for all the catalysts at 393 K. The compound labeled as “B” shows a typical profile of primary reaction product undergoing further reactions. By contrast, “C” intermediate follows the typical pattern of a secondary product, being its selectivity negligible at zero conversion and reaching a maximum for Pd and Pt/alumina catalysts and a plateau for Ru and Rh/alumina. For the case of Pd, significant selectivity for “C” is observed at zero conversion, suggesting its direct formation from the reactant. The evolution of the “C” intermediate at high conversions suggests that Ru and Rh/alumina are active for the hydrogenation of the aliphatic C–C bond, but not for the C–O bond hydrogenolysis. It should be noted that these catalysts did not show activity neither for hydrodeoxygenation nor for other polymerization reactions that had been reported in similar reactions carried out using Ru as active metal [35]. However, the experiments reported in this work have been performed at higher pressures and using largely higher catalyst/reactant ratios [35].

Concerning to the reaction scheme, results clearly indicate that the hydrogenation of the C=C bond takes place as a primary reaction in all the cases, yielding the molecule “B”. In the case of product “C” (simultaneous hydrogenation of the C=C double bond and hydrogenation of the furanic ring), the trends are not so clear. For Pd/supported catalyst, at low temperatures, the “C” compound appears at zero conversion (primary product), whereas at higher temperatures “C” is a secondary product, formed by hydrogenation of the furanic ring of the intermediate “B”. Although there are not reported studies on this specific reaction, works on the hydrogenation of conjugated systems suggest the possibility of simultaneous hydrogenation of multiple conjugated carbon–carbon double bonds. Loffreda et al. described the formation of butane as primary product in the hydrogenation of butadiene on Pt surfaces, justifying these findings on DFT studies [36].

Once “B” and “C” intermediates are formed, the next step is the ring opening, and the hydrogenation of the C=O bond. In all cases, the presence of furanic alcohols (resulting from a single hydrogenation of the C=O bond) and hydroxyl-carbonyl compounds (resulting from the ring opening) is discarded. Only the “D” intermediate, formed by the ring opening, yielding a terminal hydroxyl group, and C=O bond hydrogenation, has been detected. Since both functionalities are far enough for discarding a concerted mechanism,

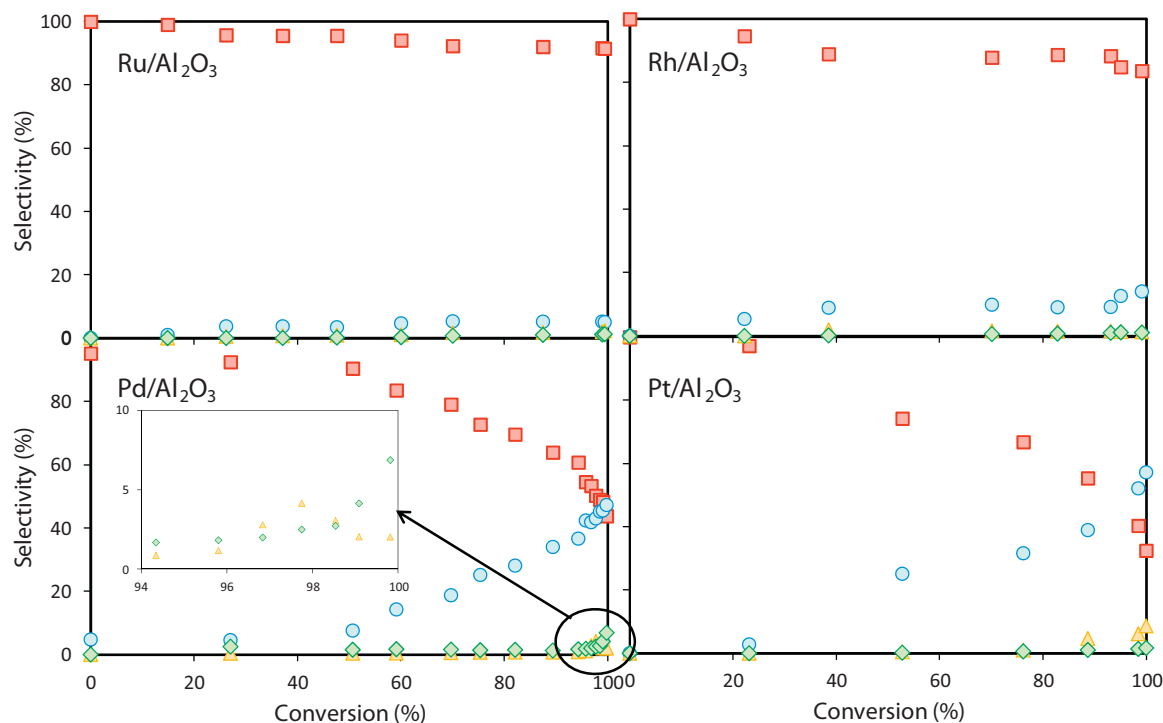


Fig. 3. Evolution of the selectivity of each intermediate as function of the C8-condensated conversion in the hydrodeoxygenation with the different catalysts used in this work (see Fig. 1 for codes).

the abovementioned non-observed intermediates are unstable and undergo fast reactions yielding the intermediate “D”. The high reactivity of the linear oxygenated intermediates is also supported by the low selectivities found for “D” intermediate, with maximum values always lower than 10%.

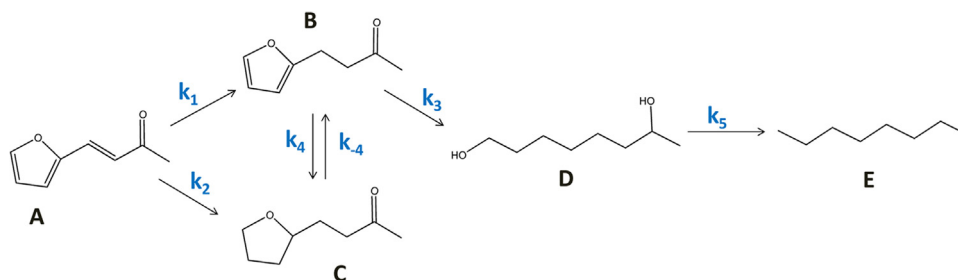
To the best of our knowledge, there are not systematic studies comparing the rates of both kinds of reactions (ring opening, and C=O bond hydrogenation), although it is suggested that both of them are slower than the hydrogenation of C=C bonds [37]. Concerning to the precursor of “D” intermediate, at first insight, ring opening product can derive either from “B”, “C” or both of them. However, *Sitthisa and Resasco* studied the ring-opening reaction of furan over Ni and Pd catalysts [38]. They observed, using both furan and tetrahydrofuran reactants, that the ring opening rates are larger with the unsaturated ring. Thus, the strongest interaction of the aromatic ring with the metal, weakening the C–O bond, plays a key role in the ring opening reaction [39]. Once the intermediate “D” is formed, it undergoes fast dehydration-hydrogenation steps yielding octane.

According to this mechanism, it is predicted that the intermediate “C” shows the behavior of a non-reactive secondary product, not undergoing further reaction. Although this is the most common pattern, there are cases (especially with Rh and Pt catalysts) in which the selectivity for the formation of this compound shows

a slight decrease at higher conversions. This behavior is explained considering that the hydrogenation of the furanic ring can be partially reversible. Equilibrium in the hydrogenation of aromatic cycles was previously observed for the hydrogenation of polyaromatic compounds [40].

Other side reactions are discarded due to the high carbon balance closures and the chemical analysis of the reaction mixture. GC–MS identification analyses were carried out, detecting only compounds with eight carbon atoms. Consequently, polymerization and hydrocycloaddition reactions were discarded at our reaction conditions. Interestingly, neither reaction products resulting from C–C bond cleavage were found. By contrast, during the hydrodeoxygenation of furfural over supported noble metals, important amounts of decarbonylation products (furan and CO) were found, specially using Pd as catalyst [41]. It is remarkable that the carbonylic carbon atom is more electropositive in the furfural molecule than in the intermediate “B”, where the terminal methyl group and the intermediate carbon chain decrease the electropositive character of this carbon atom (inductive effect). Taking into account all these facts, a kinetic model as the depicted in Scheme 3 is proposed for modeling the reaction.

The temporal evolution of the concentration of reactant, reaction intermediates and the n-octane with reaction time was modeled using the SCIENTIST® software, applying the



Scheme 3. Reaction mechanism proposed to the hydrodeoxygenation of the furfuryldeneacetone.

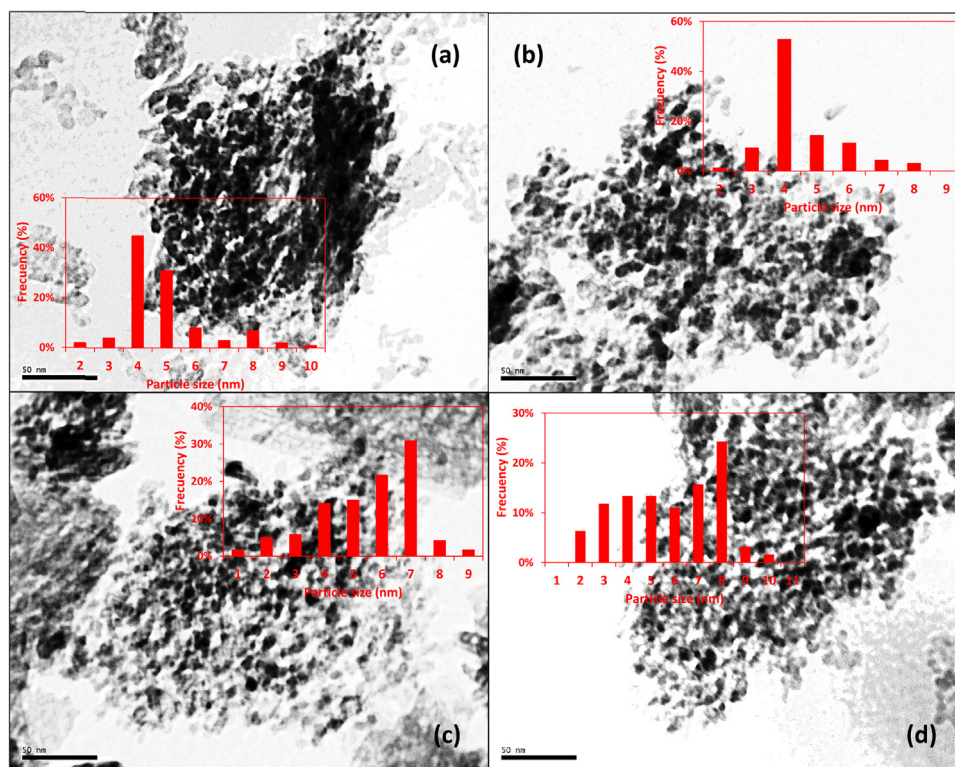


Fig. 4. TEM micrographs: (a) Ru/Al₂O₃, (b) Rh/Al₂O₃, (c) Pd/Al₂O₃ and (d) Pt/Al₂O₃.

Burlich–Stoer approximation in order to determine the kinetic constants for each step. Each elementary step has been considered to follow a pseudo-first order kinetics on the organic reactant. The presence of different organic reactants with similar molecular structure prevents the use of more mechanistic models, such as Langmuir–Hinshelwood ones. The presence of mass transfer effect has been discarded by ensuring the small catalytic particle size and with the high stirring, so this reaction is considered to be kinetically controlled. The high pressure (4.5 MPa) ensures that the hydrogen is in enough excess to consider zeroth order kinetic dependence. In any case, the negligible decrease in the pressure during the reaction leads us to neglect the effect of hydrogen pressure on the reaction kinetics at the studied interval. First-order kinetic dependence on the organic molecule has been considered for each step, which is in good agreement with most of the kinetic studies of liquid-phase hydrogenations [40,42]. Therefore, and considering batch reactor and the reaction scheme suggested in the analysis of the conversion-selectivity plots, the following set of differential equations are used for modeling the reaction:

$$\frac{dA}{dt} = -k_1 \cdot A - k_2 \cdot A \quad (I.1)$$

$$\frac{dB}{dt} = k_1 \cdot A - k_3 \cdot B - k_4 \cdot B + k_{-4} \cdot C \quad (I.2)$$

$$\frac{dC}{dt} = k_2 \cdot A + k_4 \cdot B - k_{-4} \cdot C \quad (I.3)$$

$$\frac{dD}{dt} = k_3 \cdot B - k_5 \cdot D \quad (I.4)$$

$$\frac{dE}{dt} = k_5 \cdot D \quad (I.5)$$

Different variations of these models were also tested, increasing the number of reactions considered as reversible, or including a direct transformation of intermediate “C” into intermediate “D”. In all the cases, fitting results were worse than in the proposed model. Thus, this kinetic model has been used for comparing the results obtained with the different catalysts. Likewise, the kinetic values were normalized considering the dispersion of the active phase according to the following expression and considering the characterization results summarized in Table 1.

$$k'(\text{L s}^{-1} \text{ at } -\text{g}_{\text{metal}}^{-1}) = \frac{k(\text{s}^{-1}) \cdot V(\text{L})}{M_{\text{cat}}(\text{g}) \cdot (\% \text{metal}/100) \cdot (1/A_{\text{metal}}) \cdot d}$$

where “k” is the first-order kinetic constant for a reaction rate expressed as mol reacted/formed per second and volume of feed; “k’” is the first-order kinetic constant for a reaction rate expressed as mol reacted/formed per second and exposed atom of metal; “M_{cat}” is the mass of catalyst used in each reaction; “A_{metal}” is the

Table 1

Summary of the main morphological and chemical properties (TEM, N₂ physisorption and TPD-NH₃) of the catalysts used in this work.

Catalyst	TEM		Morphological properties ^a			Acid sites (μmol/g, (K))	
	Dispersion (%)	Particle size (nm)	Specific surface area (m ² /g)	Pore diameter (nm)	Pore volume (cm ³ /g)	Weak	Medium strength
Ru-Al ₂ O ₃	23	5	118	16.8	0.50	662 (409), 335 (509)	870 (642)
Rh-Al ₂ O ₃	24	4	117	16.5	0.49	895 (408), 187 (532)	702 (636)
Pd-Al ₂ O ₃	14	8	92	18.2	0.46	600 (337), 281 (553)	610 (685)
Pt-Al ₂ O ₃	24	4	116	17.0	0.49	798 (453)	119 (660)

^a Results adapted from references Díaz et al. [29,30].

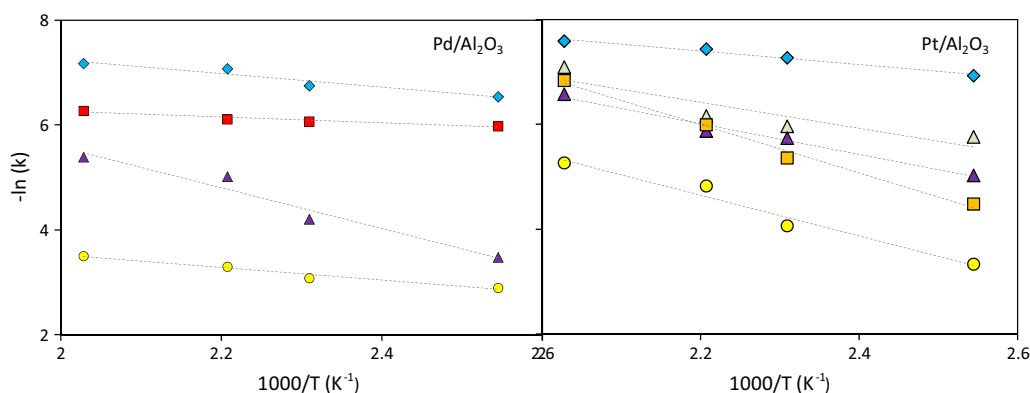
Table 2Kinetic constants ($\text{L s}^{-1} \text{at-}g_{\text{cat exp}}$) for the fitting of the experimental results at 393 K to the proposed kinetic model.

	k_1'	k_2'	k_3'	k_4'	k_5'	k_{-4}'	r^2
Pd/ Al_2O_3	9.6×10^2	1.3×10^2	2.1×10^1	–	2.8×10^2	–	0.98
Pt/ Al_2O_3	1.0×10^3	–	3.0×10^1	3.4×10^2	1.6×10^2	9.4×10^1	0.98
Rh/ Al_2O_3	4.8×10^2	–	–	6.4×10^1	–	1.8×10^2	0.98
Ru/ Al_2O_3	5.0×10^2	–	–	5.7×10^1	–	8.7×10^2	0.97

Table 3

Activation energy and pre-exponential value obtained by Arrhenius model.

	Pd/ Al_2O_3			Pt/ Al_2O_3		
	$k_{0,i}$	E_a (kJ/mol)	r^2	$k_{0,i}$	E_a (kJ/mol)	r^2
k_1	1.9×10^4	10.9	0.92	2.8×10^4	10.8	0.98
k_2	1.6×10^3	4.6	0.94	–	–	–
k_3	3.8×10^2	32.2	0.97	5.4×10^5	32.3	0.96
k_4	–	–	–	1.4×10^5	20.5	0.90
k_5	6.2×10^5	10.0	0.95	2.5×10^5	24.3	0.98
k_{-4}	–	–	–	1.0×10^7	38.3	0.99

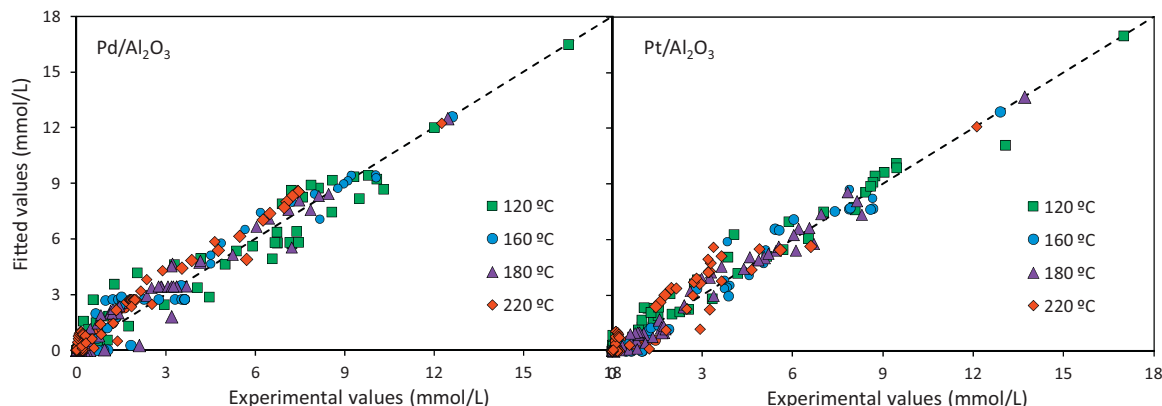
**Fig. 5.** Arrhenius dependence of the kinetic constants (see Scheme 3) on the absolute temperature for the HDO of the condensation adduct over Pd (left) and Pt (right catalysts). Kinetic constants: k_1' (◆), k_2' (■), k_3' (●), k_4' (▲), k_5' (▲), k_{-4}' (■).

atomic mass of each metal; and “d” is the metal dispersion determined by TEM.

Considering the kinetic constants normalized by exposed atom of metal, possible effects of the different effective metal loading were discarded. In any case, micrographs obtained by TEM demonstrate a high similarity in terms of particle size among the four materials, with a narrow distribution around the average value, as it can be observed in Fig. 4. Since also morphology of the catalysts and acid properties are very similar for the four studied catalysts, differences in catalytic performance are caused by the

activity of the metal. Values of the kinetic constants for the experiments performed at 393 K are reported in Table 2. The proposed model (considering the above-mentioned peculiarities for each catalyst) provides a good fit for all the experiments performed (with a correlation coefficient higher than 0.97 for all the compounds in all the reactions). The goodness of these fits is also observed in Fig. 1 for the experiments performed at 393 K with the four catalysts.

According to the proposed mechanism, Pd and Pt catalysts present similar activity for the first step of the hydrodeoxygenation (aliphatic C=C bond hydrogenation), with the highest values

**Fig. 6.** Summary of the results obtained at different temperatures in terms of parity plots for the concentration of the reactant and all the reaction products.

of k_1 . This result is in good agreement with the well-known high activity and selectivity of these metals in C=C aliphatic hydrogenations [37]. In the case of Rh and Ru catalysts, the activity is lower. These facts are in a good agreement with the trends reported for olefins hydrogenation, Pt and Pd being considered the most active catalysts for this reaction, whereas the Rh and Ru are markedly less active. Concerning to the simultaneous hydrogenation of the double bond and the furanic ring, this reaction only takes place with the Pd catalyst and at very low reaction rates (kinetic constant is ca. one order of magnitude lower than k_1). In the other cases, the intermediate “C” is only formed by hydrogenation of the intermediate “B”. In this case, the supposition of a reversible step for this transformation is supported by the values of the same order of magnitude obtained for the constants k_4 and k_{-4} . The high values of the second kinetic constant for Ru and Rh catalysts explain the slight decrease in the concentration of this intermediate, which does not undergo further transformations. At this point, the behavior of Pd differs from the other metals, being observed that the formation of intermediate “C” is irreversible.

The ring-opening reactions only take place at relevant extent with Pd and Pt catalysts (k_3), Pt being the most active catalyst, whereas Pd seems to be more active for the final dehydration-hydrogenation step (k_5). Comparing both steps, the first one is slower, in good agreement with the profile observed for the aliphatic diol “D”. It should be noted that catalyst acidity can play a key role in the dehydration-hydrogenation step, being the acid site distribution shifted to higher concentration and strength of the acid sites for the Pd catalyst.

The different behavior of the catalysts cannot be explained by the morphology or metal dispersion of these materials (Table 1) and, consequently, the activity can be attributed to the different affinity between the metal crystallites and the reactant and intermediate products. Some researches indicated that the activity in similar processes is conditioned by the interaction of unsaturated functional groups and hydrogen with the metal and the subsequent ability of the metal to eliminate the hydrogen atom [43]. The most stable interactions presented in Pd and Pt are π -allyl modes, and in the case of Pd this interaction is so strong that difficult desorption of the first intermediate (“B”). On the other hand, the hydrogenation of the aldehyde group is not favored with Ru and Rh due to their low preference of these metallic surface for C=O. This affinity is important in order to stabilize the hydrogenation intermediates in molecules with ketonic groups and it is higher in the case of Pd or Pt [44].

Kinetic values obtained by fitting the experiments catalyzed by Pd and Pt catalysts at higher temperatures (433, 453 and 493 K) are in good agreement with the similar behaviors obtained with both catalysts at low temperatures. Kinetic constants obtained using the proposed model, at different temperatures, were used for estimating the values of the pre-exponential factor and the activation energy of each step of this reaction considering an Arrhenius model. Values obtained are summarized in Table 3. Straight lines obtained are plotted in Fig. 5, observing a good linear trend. The goodness of the proposed model is also observed in Fig. 6, with a correlation coefficient higher than 0.97 for all the compounds in all the reactions. In general terms, the kinetics constants of all the relevant steps are higher and show sharper increases with temperature with Pt, in good agreement with the higher octane selectivity observed.

Although, at first insight, the values of the activation energies seem lower than expected, similar values were recently reported for HDO. Activation energies of about 10 kJ/mol are obtained for the propanoic acid HDO using supported Pd catalysts [45]. Concerning to biomass model compounds (furans, aldehydes, etc.), activation energy values ranging from 48 kJ/mol of furfural hydrodeoxygenation catalyzed by Cu/SiO₂ [16] to 70 kJ/mol of phenol HDO catalyzed by Pd/C [46] were reported. Kinetic studies of individual steps in

hydrodeoxygenation processes catalyzed by noble metals using different solvents justified the Ea lower values obtained (from 12 to 28 kJ/mol) by a strong solvation of the adducts and the metal surface by the organic solvent [46]. Although obtained values might suggest the presence of external or internal mass transfer limitations, both of them have been discarded by ensuring that particle size and stirring reaction do not affect obtained results, and also by theoretical considerations based on the Frössling equation and the Sherwood, Reynolds and Schmidt dimensionless numbers [47]. The calculated values for the reactant mass transfer coefficient ($k_g = 4.7$ m/s) and internal diffusivity (Thiele modulus value of 3.6×10^{-6}) for the most unfavorable case (Pt/Al₂O₃ at 493 K) indicate insignificant external and intraparticle mass transfer limitations under the investigated conditions.

In general terms, the activation energies of the main route reactions (A→B→D→E) are very similar for both catalysts, except in the last step whose activation energy is lower for Pd. In both cases, the lowest kinetic constant and the largest activation rates were found for the ring-opening reaction, suggesting that this is the controlling step in this reaction.

4. Conclusions

Four alumina-supported metals were tested in the hydrodeoxygenation of the C8-condensated adduct. Ru and Rh/alumina were discarded because of the negligible n-octane selectivity obtained after 8 h at 393 K. Otherwise, Pd and Pt/alumina showed high activity with selectivities to total hydrogenated products at 493 K, higher than 25 and 35%, respectively.

The activity results were used to analyze the mechanism and the catalytic properties more relevant for this reaction. The different profiles obtained with Pd and Pt catalysts were justified considering the different interaction between the in saturations and the metal surface. A kinetic model has been proposed and experimentally validated.

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